

Customer Number: 000959

DIVISIONAL-CONTINUATION APPLICATION TRANSMITTAL FORM UNDER RULE 1.53(b) (former Rule 1.60)

DOCKET NUMBER	ANTICIPATED CLASSIFICATION OF THIS APPLICATION:		PRIOR APPLICATION SERIAL NUMBER: 08/170,579	PRIOR APPLICATION FILING DATE: DECEMBER 20, 1993
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ASSISTANT COMMISSIONER FOR PATENTS **BOX PATENT APPLICATION** WASHINGTON, DC 20231

	Г	CERTIFICATION UNDER 37 CFR 1.10
	I	Date of Deposit: January 23, 1998 Mailing Label Number: EM284254988US
	b "	hereby certify that this 37 CFR 1.53(b) request and the documents referred to as attached therein are being deposited with the United States Postal Service on the date indicated above in an envelope as Express Mail Post Office to Addressee" service under 37 CFR 1.10 and addressed to the Assistant Commissioner for Patents, Box Patent Application, Washington, D.C. 20231.
	- N	Nelson Barros Name of Person Mailing Paper Signature of Person Mailing Paper
De	ar S	ir:
app	licati	request for filing a continuation divisional application under 37 CFR 1.53(b), of pending prior ion serial no. 08/170,579 filed on December 20, 1993 , of Harry Meade; Daniel Pollock; Paul entitled TRANSGENIC PRODUCTION OF ANTIBODIES IN MILK
1.		Enclosed is a copy of the latest inventor signed application, including the oath or declaration as originally filed. The copy of the enclosed papers is as follows: X
2.		A verified statement to establish small entity status under 37 CFR 1.9 and 1.27, a copy of which is enclosed, was filed in the prior application and such status is still proper and desired (37 CFR 1.28(a)).
3.	X	The filing fee is calculated below:

3.

		NUMBER OF CLAIMS FILED					IUMBER EXTRA
TOTAL	*	18	MINUS	**	20	ļ= '	0
INDEP.	*	3	MINUS	***	3	-	0
☐ MULTIPLE DEPENDENT CLAIMS							

SMALL ENTITY			
RATE FEE			
x 11 =	\$.00		
x 41 =	\$.00		
+135 =	\$.00		
BASIC FEE	\$.00		
TOTAL	\$0.00		

OTHER THAN A SMALL ENTITY RATE FEE <u>or</u> x 22 = \$0.00 \$ 0.00 x 82 =\$.00 + 270 = BASIC FEE \$790.00 <u>or</u> \$790.00 TOTAL

4.	X	The Commissioner is hereby authorized to charge any additional fees which may be required, or credit any overpayment, to Deposit Account No. 12-0080. A duplicate copy of this sheet is enclosed.				
5.	X	A check in the amount of $$990.00$ is enclosed for payment of the filing fee.				
6.		Cancel in this application original claimsof the prior application before calculating the filing fee. (At least one original independent claim must be retained for filing purposes.)				
7.		A preliminary amendment is enclosed. (Claims added by this amendment have been properly numbered consecutively beginning with the number next following the highest numbered original claims in the prior application.)				
8.	X	Amend the specification by inserting before the first line the sentences: "This application is a divisional application of serial no. 08/170,579 filed on December 20, 1993, Pending. The contents of all of the aforementioned application(s) are hereby incorporated by reference."				
9.		Please abandon said prior application as of the filing date accorded this application. A duplicate copy of this transmittal is enclosed for filing in the prior application file. (May be used if signed by person authorized by §1.138 and before payment of base issue fee.)				
10.	X	Transfer the drawings from the pending prior application to this application.				
11.		Priority of application serial nofiled oninis claimed under 35 U.S.C. §119. The certified copy has been filed in prior application serial nofiled on The certified copy will follow.				
12.	X	The prior application is assigned of record to Genzyme Transgenics Corporation.				
13.		Amonth extension of time has been submitted in the parent application Serial No. in order to establish copendency with the present application.				
14.	X	Also enclosed is/are:				
		a. A copy of the Power of Attorney from Darby & Darby P.C. to William G. Gosz, Esq. of the Genzyme Corporation.				
		b. 🖾 A copy of the Power of Attorney from William G. Gosz. Esq. of the Genzme Corporation to Lahive & Cockfield, LLP.				
		c. A copy of the Sequence Listing as filed in the prior application.				
15.	X	The power of attorney in the prior application is to <u>Lahive & Cockfield, LLP</u> .				
		a.				
		b. \square Since the power does not appear in the original papers, a copy of the power in the prior application is enclosed.				
		c. A new power has been executed and is attached.				
16.	X	Address all future communications (May only be completed by applicant, or attorney or agent of record) to Louis Myers at Customer Number: 000959 whose address is:				

Lahive & Cockfield, LLP 28 State Street Boston, Massachusetts 02109

- 17. Any requests for extensions of time necessary in a parent application for establishing copendency between this application and a parent application are hereby requested and the Commissioner is authorized to charge any fee associated with such an extension to Deposit Account No. 12-0080.
- 18. Pursuant to 37 CFR 1.821(e), the computer readable form of the sequence listing for this new application is to be identical with the computer readable form of application serial no. 08/170,579 . Please use the computer readable form of application serial no. 08/170,579 in lieu of filing a duplicate computer readable form in this application. Pursuant to 37 CFR 1.821(f), the content of the paper copy of the sequence listing for this new application and the computer readable form of application serial no. 08/170,579 are the same.

January 23, 1998

Date

LAHIVE & COCKFIELD, LLP 28 State Street Boston, Massachusetts 02109 Tel. (617) 227-7400

Louis Myers

Reg. No. 35,965

- ☐ inventor(s) ☐ filed under §1.34(a)
- ☐ assignee of complete interest
- attorney or agent of record

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4444/08981

TRANSGENIC PRODUCTION OF ANTIBODIES IN MILK

Field of the Inventi m

This invention pertains to a method for the production of monoclonal antibodics in mammal's milk, specifically through the creation of transge ic animals that selectively express foreign antibody genes in mammary epithelial cells.

Background of the In vention

Immunoglobulins are heteropolymeric proteins that are normally synthesized, modified, assembled, and secreted from circulating B lymphocytes. Using recombinant DNA technology, it is possible to program cells other than B-lymphocytes to express The difficulties encountered in this effort immunoglobulin genes 1) Both heavy and light chains of stem from several factors: immunoglobulins must be co-expressed at appropriate levels; 2) Nascent immunoglobul n polypeptides undergo a variety of co- and post-translational medifications that may not occur with sufficient fidelity or efficiency in heterologous cells; 3) Immunoglobulins require accessory chaperone proteins for their assembly; 4) The synthetic and secretory capacity of the cell may be inadequate to secrete large amounts of heterologous proteins; and 5) The secreted immunoglobulins may be unstable in the extracellular milieu of a foreign cell.

Because immunoglobulins have many therapeutic, diagnostic and industrial applications, there is a need in the art for expression systems n which these proteins can be reproducibly manufactured at a high level, in a functional configuration, and in

a form that allows them to be easily harvested and purified. The development of transgenic animal technology

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has raised the possibility of using large animals as genetically P.C.T. application WO 90/04036 programmed protein factories. (published 4/19/90) discloses the use of transgenic technology for immunoglobulin expression. WO 92/03918 (3/19/92) and WO 93/12227 (6/24/93) teach the introduction of unrearranged immunoglobulin genes into the germline of transgenic animals. The use of intact immunoglobulin genes (including their respective promoter regions) will result in their expression in lymphocytes and secretion intothe bloodstream of the host animal; this necessitates a strategy. endogenous host's expression the of the for suppressing purifying of the problem immunoglobulins, raises and immunoglobulins from serum, which contains many other proteins, including proteolytic enzymes. Furthermore, if the transgenic approach is chosen, heavy and light chain genes must both be incorporated into the host genome, in a manner that enables their comcomittant expression.

Another option in creating transgenic animals is to link the gene of interest to a heterologous transcriptional promoter that only functions in a defined cell type within the host. this manner, tissue-specific expression of the transgene may be U.S. Patent No. 4,873,316 (issued October 10, 1989) programmed. the production of recombinant tissue plasminogen discloses activator (TPA) in the milk of transgenic mice in which the TPA gene is linked to the promoter of the milk protein casein. proteins that have been expressed in a similar fashion include cystic fibrosis transmembrane conductance regulator (DiTullio et Bio/Technology 10:74, 1992), urokinase (Meade al., 1990), interleukin-2 (Buhler et Bio/Technology 8: 443, Bio/Technology 8:140, 1990), and antihemophilic factor IX (Clark et al., Bio/Technology 7:487, 1989). Notably, these proteins are all not require do polypeptides that simple single-chain multimerization or assembly prior to secretion.

It has now been found that when a transgenic mammal is created carrying paired immunoglobulin light and heavy chain genes under the control of the casein promoter, such an animal produces large amounts of assembled immunoglobulins which are secreted in its milk. Using the DNA constructs of the present invention, a surprisingly high efficiency of co-integration of heavy and light chain genes is observed. Using the metod and constructs of the present invention, it is possible for the first time to program a mammary epithelial cell to produce and assemble complex tetrameric glycoproteins and secrete them in high quantities.

Accordingly, it is an object of the present invention to provide methods for the large-scale production of immunoglobulins in the milk of transgenic mammals.

Another object of the invention is to provide methods for the design of synthetic immunoglobulins that can be produced in large quantities in milk.

Yet another object of the invention is to provide methods for administering therapeutically beneficial antibodies to suckling young, by creating female mammals that excrete such antibodies into their milk.

A further object of the invention is a transgenic non-human mammal having germ and somatic cells with recombinant DNA sequences encoding immunoglobulin light and heavy chains, where said sequences are operatively linked at their 5' termini to a mammary specific promoter and at their 3' end to a sequence comprising a polyadenylation site.

A further object of the invention is a casein promoter cassette comprising in the 5' to 3' direction:

- a) 5' promoter sequences from the beta casein gene,
- L) an XhoI restriction site, and
- c) 3' untranslated sequences from the goat beta casein gene.

These and other objects of the present invention will be apparent to those of ordinary skill in the art in light of the present specification, drawings, and claims.

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Brief Description of the Drawings

Figure 1 is a schematic representation of the Bc62 plasmid, which contains a 13.9 kb Sal I fragment that comprises cDNA encoding immunoglobulin light chain, flanked on its 5' and 3' termini by goat beta casein sequences.

Figure 2 is a schematic representation of the Bc61 plasmid, which contains a 14.6 kb Sal I fragment that comprises cDNA encoding immunoglobulin heavy chain, flanked on its 5' and 3' termini by goat beta casein sequences.

Figure 3 depicts the immunoblot detection of human immunoglobulin heavy chain in the milk of transgenic mice that were created using the beta casein promoter-linked immunoglobulin genes shown in Figures 1 and 2.

Figure 4 depicts the immunoblot detection of human immunoglobulin light chain in the milk of transgenic mice that were created using the beta casein promoter-linked immunoglobulin genes shown in Figures 1 and 2.

Summary of the Invention

In one aspect, this invention comprises a method for obtaining heterologous immunoglobulins from the milk of transgenic mammals. Another aspect of the prevent invention comprises the method for creating transgenic mammals by introducing into their germline immunoglobulin cDNA linked to a milk-specific promoter.

In another aspect, the present invention comprises transgenic mammals having germ cells and somatic cells having recombinant DNA sequences comprising immunoglobulin cDNA linked to a milk-specific promoter.

In still another aspect, the present invention comprises an isolated DNA comprising an expression cassette having 5' and 3' non-coding sequences derived from the goat beta casein gene linked via a unique restriction site that serves as a convenient cloning site for immunoglobulin coding sequences.

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Detailed Description of the Invention

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All patent applications, patents and literature cited in this specification are hereby incorporated by reference in their entirety. In the case of inconsistencies, the present disclosure will prevail.

The present invention pertains to a method for the production of monoclonal antibodies that are excreted into the milk of transgenic animals and the method for production of such animals. This is achieved by engineering DNA constructs in which DNA segments encoding specific paired immunoglobulin heavy and light chains are cloned downstream of a promoter sequence that is preferentially expressed in mammary epithelial cells. recombinant DNAs containing the promoter-linked heavy and light chain genes are then coinjected into preimplantation embryos. progeny are screened for the presence of both transgenes. Representative females from these lines are then milked, and the milk is analyzed for the presence of the monoclonal antibody. order for the antibody to be present, both heavy and light chain genes must be expressed concurrently in the same cell. antibodies may be purified from the milk, or the milk itself, comprising the immunoglobulins, may be used to deliver antibodies to a recipient. This is discussed below.

The immunoglobulin genes useful in the present invention may be obtained from natural sources e.g. individual B cell clones or hybridomas derived therefrom. Alternately, they may comprise synthetic single-chain antibodies in which the light and heavy variable regions are expressed as part of a single polypeptide. Furthermore, recombinant antibody genes may be used that have been predictively altered by nucleotide substitutions that do or do not change the amino acid sequence, by addition or deletion of sequences, or by creation of hybrid genes in which different regions of the polypeptide are derived from different sources. Antibody genes by their nature are extremely diverse, and thus naturally tolerate a great deal of variation. It will be appreciated by those skilled in the art that the only limitation

for producing an antibody by the method of the present invention is that it must assemble into a functional configuration and be secreted in a stable form into the milk.

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The transcriptional promoters useful in practicing the present invention are those promoters that are preferentially activated in mammary epithelial cells, including promoters that control the genes encoding milk proteins such as caseins, beta lactoglobulin (Clark et al., (1989) Bio/Technology 7: 487-492), whey acid protein (Gordon et al., (1987) Bio/Technology 5: 1183-1187), and lactalbumin (Soulier et al., (1992) FEBS Letts. 297: 13). Casein promoters may be derived from the alpha, beta, or kappa casein genes of any mammalian species; a preferred promoter is derived from the goat beta casein gene (DiTullio, (1992) Bio/Technology 10:74-77).

invention, a unique XhoI in the present restriction site is introduced at the 3' terminus of the promoter sequence to allow the routine insertion of immunoglobulin coding sequences. Preferably, the inserted immunoglobulin gene is flanked on its 3' side by cognate genomic sequences from a mammary-specific gene, to provide a polyadenylation site and transcript-stabilizing sequences. Transcription of the construct in vivo results in the containing casein-derived stable mRNA production of а untranslated sequences upstream of the translational initiator codon of the immunoglobulin gene and 3' untranslated sequences translational termination codon the downstream of immunoglobulin gene. Finally, the entire cassette (i.e. promoterimmunoglobulin-3' region) is flanked by restriction sites that enable the promoter-cDNA cassette to be easily excised as a single This facilitates the removal of unwanted prokaryotic vector-derived DNA sequences prior to injection into fertilized eggs.

The promoter-linked immunoglobulin heavy and light chain DNAs are then introduced into the germ line of a mammal e.g. cow, sheep, goat, mouse, oxen, camel or pig. Mammals are defined herein as all animals, excluding humans, that have mammary glands and produce milk. Mammalian species that produce milk in large amounts over long periods of time are preferred. Typically, the DNA is injected into the pronuclei of fertilized eggs, which are then implanted into the uterus of a recipient female and allowed to gestate. After birth, the putative transgenic animals are tested for the presence of the introduced DNA. This is easily achieved by Southern blot hybridization of DNA extracted from blood cells or other available tissue, using as a probe a segment of the injected gene that shows no cross hybridization with the DNA of the recipient species. Progeny that show evidence of at least one copy of both heavy and light-chain immunoglobulin genes are selected for further analysis.

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Transgenic females may be tested for immunoglobulin secretion into milk, using any of the immunological techniques that are standard in the art (e.g. Western blot, radioimmunoassay, ELISA). The anti-immunoglobulin antibodies used in this analysis may be polyclonal or monoclonal antibodies that detect isolated heavy or light chains or others that react only with fully assembled (H2L2) immunoglobulins.

The recombinant immunoglobulins are also characterized with respect to their functionality, i.e. binding specificity and affinity for a particular antigen. This is achieved using immunological methods that are standard in the art, such as Scatchard analysis, binding to immobilized antigen, etc. The stability characteristics of an immunoglobulin in the milk of a given species are also assayed, by applying the above-described detection methods to milk that has been incubated for increasing times after recovery from the animal.

The immunoglobulins produced by the methods of the present invention may be purified from milk, using adsorption to immobilized Protein G, column chromatography, and other methods known to those of ordinary skill in the art of antibody purification.

The level of production of recombinant immunoglobulins in an individual transgenic mammal is primarily determined by the site

and manner of integration of the transgene after injection into the fertilized egg. Thus, transgenic progeny derived from different injected eggs may vary with respect to this parameter. The amount of recombinant immunoglobulin in milk is therefore monitored in representative progeny, and the highest-producing females are preferred.

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Those skilled in the art will recognize that the methods of the present invention can be used to optimize the production of natural and synthetic immunoglobulins. The steps of creating a transgenic animal, testing for the presence of both heavy and light-chain genes, assaying the secretion of immunoglobulin into the milk of female progeny, and, finally, assessing the quality of the resulting antibodies, can be repeated sequentially, without undue experimentation, to establish preferred constructs for different applications.

According to the present invention, the nature of the recombinant immunoglobulins and their specific mode of use can vary. In one embodiment, the present invention encompasses high-level expression of antibodies that are harvested and purified from milk and used in purified form. High-level expression is defined herein as the production of about 1 mg/ml of protein. In another embodiment, antibodies are engineered that provide protection to humans against infectious diseases; therapeutic administration is then achieved by drinking the milk. In a still further embodiment, lactating animals are engineered to produce antibodies specifically beneficial to their offspring, which acquire them through suckling. In a still further embodiment, animals produce an antibody that protects the lactating mammal itself against breast pathogens e.g. bacteria that produce mastitis.

The unexpectedly high-volume expression of immunoglobulins using the method and constructs of the present invention also allows the use of such immunoglobulins in pharmaceutical and chemical settings. By way of non-limiting example the method of the present invention can be used to produce high levels of tetrameric antibodies directed against various

Salmonella, hepatitis B coli, (e.g. E. pathogens erythropoietin, tissue (e.g. active peptides biologically plasminogen activator, gamma interferon) and for use in chemical reactions directed against various enzymes. Monoclonal antibodies that bind to the transition state of a chemical reaction can be Furthermore, monoclonal used in industrial-scale production. antibodies are often immobilized on columns for use in the purification of biopharmaceuticals; in such cases, production of the antibodies represents a significant fraction of the cost of purification. The methods of the present invention facilitate the production of high-volume, low cost antibody stocks for use in these types of applications.

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The present invention is further described following working examples, which are intended to illustrate the invention without limiting its scope.

Construction of a Milk-Specific Promoter Cassette Example 1:

The present invention encompasses a recipient vector into which many different immunoglobulin genes can be interchangeably inserted. The vector contains 5' milk-specific promoter sequences and 3' untranslated genomic sequences that flank an XhoI cloning site. This cloning is unique because it is the only one present in the vector. Preferably, the entire expression cassette should be flanked by restriction sites that allow the easy excision of the promoter-linked immunoglobulin gene.

In this Example, the promoter and 3' genomic sequences were derived from the goat beta casein gene. The gene was cloned and characterized as described by Roberts et al., 1992, Gene 121:255-262, which is hereby incorporated by reference.

of insertion prior The expression cassette, upstream of consists of 6.2 kb immunoglobulin genes, translational start of the beta casein coding sequence and 7.1 kb of genomic sequence downstream of the translational stop of the beta casein gene. The TaqI site just upstream of the translational start codon was changed to an XhoI site. This unique XhoI cloning site is at the junction of the upstream and downstream sequences. It is this XhoI site, included in the sequence CGCGGATCCTCGAGGACC, into which recombinant immunoglobulin genes are inserted. Bio/Technology 10:74-77) Tullio, (1992)

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The 3' beta casein region begins at the PpuMI site found Included in this in Exon 7 and continues for 7.1 kb downstream. sequence are the remaining 18 bp of Exon 7, and all of Exon 8 and These encode the 3' untranslated regions of the goat beta Exon 9. with the sequence: terminate casein gene, anđ

To engineer restriction, sites, flanking the casein cassette, the goat beta casein control sequences were first cloned into the SuperCosl vector (#251301, Stratagene, La Jolla, CA) with flanking NotI and SaII sites. This plasmid was then modified by This created a 13.3 kb changing the NotI site to a SaII site. SalI fragment containing the beta casein expression cassette within the qbc163 vector.

Example 2: Construction of Promoter-linked Monclonal Antibody Genes

In this Example, the genes encoding a human monoclonal antibody directed against a colon cancer cell-surface marker were linked to the casein promoter. cDNAs encoding the light and heavy chains of this antibody were cloned from an antibody-secreting hybridoma cell line into a pUC19-derived vector. heavy chain cDNAs were present on HindIII/EcoRI fragments of 702 bp and 1416 bp, respectively.

To adapt the genes for insertion into the casein promoter cassette, XhoI restriction sites were engineered at both ends of each DNA segment as detailed below. In the same step, the region upstream of the immunoglobulin translation initiation codon was modified so that it contained sequences similar to those in the analogous region of the beta casein gene.

Light chain gene: The pUC19 plasmid containing the light chain cDNA insert was digested with HindIII, blunt-ended by treatment with the Klenow fragment of DNA Polymerase I, and ligated

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to an oligonucleotide containing an XhoI recognition sequence (#1030, New England Biolabs, Beverly, MA).

The region immediately upstream of the initiating ATG was then mutagenized using an oligonucleotide with the following sequence: 5' AGT GAA TTC ATG CTC GAG AGC CAT GGC CTG GATC 3'. Digestion of the final plasmid with XhoI produced the modified light chain cDNA that was flanked by Xhol cohesive ends.

The light chain cDNA was then inserted into the unique XhoI cloning site of the gbcl63 expression vector described in Example 1, yielding plasmid Bc62 (Figure 1).

chain cDNA was mutagenized using an oligonucleotide with the following sequence: 5' AGT GAA TTC ATG CTC GAG AGC CAT GAA GCA CCTG
3'. The resulting plasmid contains an XhoI site upstream of the heavy chain translation initiation codon.

The downstream HindIII site was converted to an Xhol site using a synthetic adapter with the sequence 5' AGC TCC TCG AGG CC 3'. Digestion of the modified plasmid with XhoI produced the the 1.4 kb modified heavy chain cDNA flanked by XhoI cohesive ends. This fragment was then inserted into the unique XhoI cloning site of gbc163 to yield Bc61 (Figure 2).

Prior to injection, promoter-linked light and heavy chain genes were isolated from Bc61 and Bc62, respectively, by digestion with SalI. The fragments were then purified by gel electrophoresis followed by CsCl equilibrium gradient centrifugation. The DNA was dialyzed extensively against distilled water prior to quantitation.

Example 3: Production of Transgenic Mice

The casein promoter-linked DNA fragments encoding the immunoglobulin heavy and light chains, obtained as described in Example 2, were injected into fertilized mouse eggs using procedures that are standard in the art, as described in Hogan, B., Constantini, F., and Lacey, E., Manipulating the Mouse Embryo: A Laboratory Manual (Cold Spring Harbor Laboratories, 1986). The resulting progeny were then analyzed for the presence of both

antibody gene sequences. DNA was extracted from tail biopsy material and probed using Southern blot analysis. The probes used in the hybridization were the original cDNAs encoding the heavy and light chains. As seen in Table 1, most of the first generation transgenic progeny had incorporated both transgenes.

<u>Table 1</u>
<u>Summary of Bc61 - Bc62 Mice</u>

			· · · · · · · · · · · · · · · · · · ·	
Founder	Sex	. Bc61	Bc62 ·	Expression
1-2	M	· Pos: `	· Pos: "^	and the second of the second o
1-3	M	Pos.	Pos.	light chain only
1-9	M	Pos.	Pos.	
1-15	F	Neg.	Pos.	Low level lambda chain
1-16	F	Pos.	Neg.	
1-19	F	Pos.	Pos.	N.D.
1-23	F	Pos.	Pos.	1-3 mg/ml
1-24	F	Pos.	Pos.	low level
1-25	M	Pos.	Neg.	
1-39	M	Pos.	Pos.	
1-13	F	Pos.	Pos.	N.D.
1-56	F	Pos.	Pos.	N.D.
1-64 2-76 2-82	M F F	Pos. Pos. Pos.	Pos. Pos. Pos.	1-3 mg/ml 1-3 mg/ml
1-72 2-92 2-95	M F F	Pos. Pos. Pos.	Pos. Pos. Pos.	0.2 - 0.5 mg/ml 0.2 - 0.5 mg/ml

N.D. = not detected

Example 4: Analysis of Recombinant Immunoglobulins in Milk

Samples of milk from the transgenic mice obtained as described in Example 3 were analyzed for the presence of the

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heterologous immunoglobulin by Western blot. The heavy chain of the antibody was detected using a horseradish peroxide-linked polyclonal antibody directed against human gamma heavy chain (Antibody #62-8420, Zymed, South San Francisco, CA.) as shown in Figure 3. The light chain was detected using antibodies to the human lambda light chain, (Antibody #05-4120, Zymed, South San Francisco, CA) shown in Figure 4. In these Figures, it can be seen that immunoreactive heavy and light chains can be detected in the milk of several animals, but not in the negative control animal CD-1. Human immunolglobulin can be detected in milk from founder 1-23 and from the progeny of the 1-76 and 1-72 founders. These animals are the second-generation females, 2-76, 2-82, 2-92, and 2-95. The levels of expression range between 0.2 mg/ml to over 1 mg/ml (Table 1).

What is claimed is:

- 1 1. A method for obtaining heterologous immunoglobulin
- 2 from the milk of a transgenic mammal comprising the steps of:
- a. introducing into the germline of said mammal
- 4 DNA comprising the protein-coding sequences of said
- 5 immunoglobulin, said DNA operatively linked at its 5' terminus to
- 6 a promoter sequence that supports the preferential expression of
- 7 said genes in mammary gland epithelial cells, and said DNA
- 8 operatively linked at its 3' terminus to a sequence containing a
- 9 polyadenylation site, and
- b. obtaining milk from said mammal.
 - 1 2. The method of claim 1 wherein said mammal is 2 selected from the group consisting of mice, cows, sheep, goats,
 - oxen, camels, and pigs.
 - 1 3. The method of claim 1 wherein said promoter is
 - 2 selected from the group consisting of the casein promoter, the
 - 3 beta lactoglobulin promoter, the whey acid protein promoter, and
 - 4 the lactalbumin promoter.
 - 1 4. The method of claim 1 wherein said immunoglobulin
 - 2 comprises heavy and light chains.
 - 1 5. The method of claim 1 wherein said immunoglobulin
 - 2 comprises a single polypeptide chain.
 - 1 6. The method of claim 1 wherein said immunoglobulin
 - 2 is of human origin.
 - 1 7. The method of claim 1 wherein said immunoglobulin
 - 2 is purified from the milk of said mammal.
 - 1 8. A transgenic non-human mammal all of whose germ
 - 2 cells and somatic cells contain recombinant DNA sequences

- 3 encoding immunoglobulin heavy and light chains, wherein said
- 4 sequences are operatively linked at their 5' termini to a
- 5 promoter sequence that supports the preferential expression of
- 6 said genes in mammary gland epithelial cells, and operatively
- 7 linked at their 3' termini to a sequence containing a
- 8 polyadenylation site.
- 9. The transgenic mammal of claim 8 wherein said
- 2 mammal is selected from the group consisting of mice, cows,
- 3 sheep, goats, oxen, camels, and pigs.

1 10. The transgenic mammal of claim 8 wherein said

and the first section of the section of

- 2 promoter is selected from the group consisting of the casein
- 3 promoter, the beta lactoglobulin promoter, the whey acid protein
- 4 promoter, and the lactalbumin promoter.
- 1 11. The transgenic mammal of claim 8 wherein said
- 2 immunoglobulin comprises heavy and light chains.
- 1 12. The transgenic mammal of claim 8 wherein said
- 2 immunoglobulin comprises a single polypeptide chain.
- 1 13. The transgenic mammal of claim 8 wherein said
- 2 immunoglobulin is of human origin.
- 1 14. An isolated purified DNA comprising in the 5' to
- 2 3' direction
- a) 5' promoter sequences from the beta casein
- 4 gene,
- b) a unique Xho I restriction site, and
- 6 c) 3' untranslated sequences from the goat beta
- 7 casein gene, wherein a) comprises nucleotides -6168 to -1 of the
- 8 goat beta casein, wherein nucleotide 1 is the first nucleotide of
- 9 the beta casein translation initation codon, b) comprises the
- 10 sequence CGCGGATCCTCGAGGACC, and c) comprises the sequence

- 11 starting at the PpuMI site found at bp648 of the beta casein cDNA
- 12 sequence, and continuing for 7.1 kb downstream,
- 13 termininating in the sequence
- 14 TAAGGTCCAGAGACCGAGACCCACTCACTAGGCAACTGGTCCGRCCAGCTGTTAAGTGA.
 - 1 15. The DNA of claim 14 wherein an immunoglobulin cDNA
- 2 is inserted into b), said DNA directing the mammary-gland-
- 3 specific expression of said immunoglobulin in transgenic animals.
- 1 16. The DNA of claim 15 wherein said immunoglobulin
- 1 comprises heavy and light chains.
- 1 17. The DNA of claim 15 wherein said immunoglobulin
- 2 comprises a single polypeptide chain.
- 1 18. The DNA of claim 15 wherein said immunoglobulin is
- 2 of human origin.

ABSTRACT

A method for the production of monoclonal antibodies in mammal's milk, through the creation of transgenic animals that selectively express foreign antibody genes in mammary epithelial cells.

SEQUENCE LISTING

i	(7)	מידואידוי) /	NΥ	INFORMATION:
1	1	nanat) i	ALI	TINE ORMALION:

- (i) APPLICANT: Meade, Harry Pollock, Daniel DiTullio, Paul
- (ii) TITLE OF INVENTION: Transgenic Production of Antibodies in Milk

and the contract of the contra

- (iii) NUMBER OF SEQUENCES: 5
- (iv) CORRESPONDENCE ADDRESS: (A) ADDRESSEE: Darby & Darby PC
 - (B) STREET: 805 Third Avenue.
 - (C) CITY: New York
- - (E) COUNTRY: US
 - (F) ZIP: 10022
 - (v) COMPUTER READABLE FORM:
 - (A) MEDIUM TYPE: Floppy disk
 - (B) COMPUTER: IBM PC compatible
 - (C) OPERATING SYSTEM: PC-DOS/MS-DOS
 - (D) SOFTWARE: PatentIn Release #1.0, Version #1.25
 - (vi) CURRENT APPLICATION DATA:

- (A) APPLICATION NUMBER: US 08/170579
- (B) FILING DATE: 20-DEC-1993
- (C) CLASSIFICATION:
- (viii) ATTORNEY/AGENT INFORMATION:
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 - (C) REFERENCE/DOCKET NUMBER: 4444/08981
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 - (C) TELEX: 236687
- (2) INFORMATION FOR SEQ ID NO:1:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 18 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: cDNA to mRNA

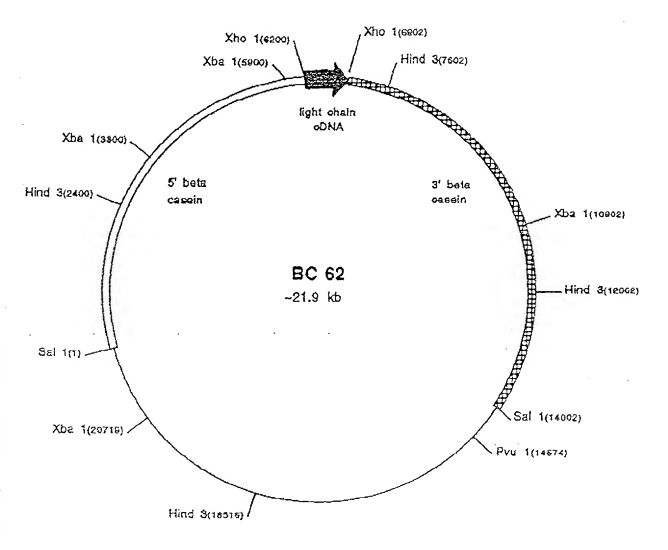
(vi) ORIGINAL SOURCE: (A) ORGANISM: Capra hircus	
(vii) IMMEDIATE SOURCE: (B) CLONE: beta casein 5'	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:1:	
CGCGGATCCT CGAGGACC	18
(2) INFORMATION FOR SEQ ID NO:2:	an garanta sa
 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 59 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 	garan Angrik di kang j
(ii) MOLECULE TYPE: cDNA to mRNA	
(vi) ORIGINAL SOURCE: (A) ORGANISM: Capra hircus	
(vii) IMMEDIATE SOURCE: (B) CLONE: beta casein 3	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:2:	
TAAGGTCCAC AGACCGAGAC CCACTCACTA GGCAACTGGT CCGTCCAGCT GTTAAGTGA	59
(2) INFORMATION FOR SEQ ID NO:3:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 34 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: cDNA to mRNA	
(vi) ORIGINAL SOURCE: (A) ORGANISM: Capra hircus	
<pre>(vii) IMMEDIATE SOURCE: (B) CLONE: light chain 5'</pre>	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:3:	
AGTGAATTCA TGCTCGAGAG CCATGGCCTG GATC	34
(2) INFORMATION FOR SEQ ID NO:4:	

(ii) MOLECULE TYPE: cDNA to mRNA
(vi) ORIGINAL SOURCE: (A) ORGANISM: Capra hircus
(vii) IMMEDIATE SOURCE: (B) CLONE: Heavy chain 5'
(XI) SEQUENCE DESCRIPTION: SEQ ID NO:4:
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	FORMATION FOR SEQ ID NO:5:
(i	(A) LENGTH: 14 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear
T (ii) MOLECULE TYPE: cDNA to mRNA
(vi	(A) ORGANISM: Capra hircus
1 :	i) IMMEDIATE SOURCE: (B) CLONE: HEAVY CHAIN 3'
	i) SEQUENCE DESCRIPTION: SEQ ID NO:5:
AGCTCC	rcga ggcc

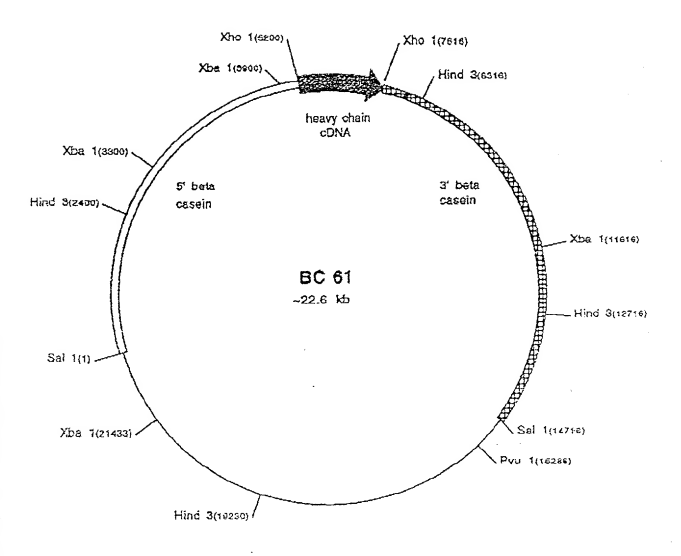
(i) SEQUENCE CHARACTERISTICS:

(D) TOPOLOGY: linear

(A) LENGTH: 34 base pairs
(B) TYPE: nucleic acid
(C) STRANDEDNESS: single



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12-76 B7 12-82 D7 12-15 D7 12-92 D7

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ALL FOREIGN APPLICATIONS, IF ANY, FILED MORE THAN 12 MONTHS PRIOR TO THE FILING DATE OF THIS APPLICATION

COUNTRY

APPLICATION NO.

DATE OF FILING

PRIORITY
CLAIMED UNDER
35 U.S.C. §119

Yes

No

POWER OF ATTORNEY:

As a named inventor, I hereby appoint the following atternoy(s) and/or agents(s) to prosecute this application and transact all business in the Patent and Trademark office connected therawith. Morris Relean #15,108, Gordan D. Copisin #19,165, William F. Dudine, Jr. #20,569, Mickael J. Sweedler #19,037, S. Peter Ludwig #25,351, Paul Fields #20,298, Joseph B. Lerch #28,838, Melvin C. Gerner #28,272, Ethan Horwitz #27,848, Baverly B. Goodwin #28,417, Adda C. Gogoris #28,714, Martin E. Goldstein #20,889, Bert J. Lewen #18,407, Hearly Sternberg #22,408, Peter C. Schechter #31,682, Rebert Schaffer #31,194, David B. Francescant #25,159

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2P 000E 020E3

I further declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United States Code, and that such willful false statements may jeopardize the validity of the application or any patent issuing thereon.

SIGNATURE OF INVENTOR 1: Harry Meade

DATED: 3/1/94

SIGNATURE OF INVENTOR 3: Paul a. D. Tallo DATED: 3/1/94

(040Fema(PTG-21)

REV. 12/87

5088729080;# 5

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Filed:

December 20, 1993

Examiner:

B. Campell

Title:

TRANSGENIC PRODUCTION OF ANTIBODIES IN MILK

I hereby certify that this correspondence is being deposited with the United States Postal Service as first class mail, postage prepaid in an envelope addressed to: The Honorable Commissioner of Patents and Trademarks, Washington, DC 20231 on the

date set forth below: April 8, 1996

Date of Signature and of Mail Deposit

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The Commissioner of Patent and Trademarks Washington, DC 20231

POWER OF ATTORNEY BY ASSIGNEE OF ENTIRE INTEREST (REVOCATION OF PRIOR POWERS)

As Assignee of record of the entire interest of the above-identified application,

REVOCATION OF PRIOR POWERS OF ATTORNEY

all powers of attorney previously given are hereby revoked and

NEW POWER OF ATTORNEY

the following attorneys are hereby appointed to prosecute and transact all business in the Patent and Trademark Office connected therewith.

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William G. Gosz	27,787
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Jennifer A. Tegfeldt	31,310
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Recorded in the PTO on December 20, 1993

Reel: 7406 Frame: 0113

4/3/96

Data

James A. Geraghty

President and CEO

Genzyme Transgenics Corporation

IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

In re the application of: Meade et al.

Serial No.: 08/170,579

Filed: December 20, 1993

For: TRANSGENIC PRODUCTION OF

ANTIBODIES IN MILK

Attorney Docket No.: 444408981

Assistant Commissioner for Patents Washington, D.C. 20231

Group Art Unit: 1804

Examiner: Campell, B.

ASSOCIATE POWER OF ATTORNEY

Sir:

The undersigned attorney for Applicant hereby appoints the following attorneys with power to prosecute and transact all business in the Patent and Trademark Office connected with the above-identified patent application:

John A. Lahive, Jr. W. Hugo Liepmann James E. Cockfield Thomas V. Smurzynski Ralph A. Loren Thomas J. Engellenner Giulio A. DeConti, Jr. Ann Lamport Hammitte Paul Louis Myers Michael I. Falkoff John V. Bianco	Reg. No. 19,788 Reg. No. 20,407 Reg. No. 19,162 Reg. No. 24,798 Reg. No. 29,325 Reg. No. 28,711 Reg. No. 31,503 Reg. No. 34,858 Reg. No. 35,965 Reg. No. 30,833 Reg. No. 36,748	Jeremiah Lynch Amy E. Mandragouras Elizabeth A. Hanley Anthony A. Laurentano Jane E. Remillard Mark A. Kurisko Beth E. Arnold Jean M. Silveri Matthew P. Vincent Lawrence E. Monks	Reg. No. 17,425 Reg. No. 36,207 Reg. No. 33,505 Reg. No. 38,220 Reg. No. 38,872 Reg. No. 38,944 Reg. No. 35,430 Reg. No. 39,030 Reg. No. 36,709 Reg. No. 34,224
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Respectfully submitted,

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Dated: 9/13/96